REMARKS/ARGUMENTS

Claims 1-4, 7, 9-19 and 41-44 are pending in this Application. The Office Action mailed on September 18, 2006, includes the following rejections:

- Claims 1-4, 7, 9-19 and 41-44 are rejected under 35 U.S.C. § 112 first paragraph.
- 2. Claims 1-4, 7, 9-19 and 41-44 are rejected under 35 U.S.C. § 112 second paragraph.
- Claims 15-17 and 19 are rejected under 35 U.S.C. § 102(b) as being anticipated by Livesey, et al.
- 4. Claims 15-19 are rejected under 35 U.S.C. § 102(b) as being anticipated by Dennis, et al.
- 5. Claims 15-19 are rejected under 35 U.S.C. § 102(b) as being anticipated by Gulati, et al.
- Claims 15-19 are rejected under 35 U.S.C. § 102(e) as being anticipated by Tanagho, et al.
- 7. Claims 15-19 are rejected under 35 U.S.C. § 102(e) as being anticipated by Atala.
- 8. Claims 1-3, 9-14 and 17 are rejected under 35 U.S.C. § 103.
- 9. Claims 4, 7 and 18 are rejected under 35 U.S.C. § 103.

Applicants respectfully address the basis for each of the Action's rejections below.

Support for the amendments to the claims can be found throughout the application. Evidence of the reduced immunogenic response may be found throughout the application. The specification supports the amendments to the claim 1, specifically paragraph [0022], which compares the acellular replacement tissue of the present invention to an allograft and shows a significantly reduced immunologic response because surface cell antigens have been removed. Paragraphs [0037-0051] state the immune response of tissue prepared with the method of the present invention show that the native cell-free tissue adapts to its environment and is not rejected. Furthermore, the composition of the present invention (the native cell-free tissue) is not rejected as other tissue replacements or allografts. The specification evaluates the native-cell free tissues for immune response following implantation of a cell-free sciatic nerve graft under various conditions, e.g., Figures 2-3 and Table 1 of the present application. In addition, paragraph [0043] compares the present invention to the current clinical approach (i.e., the autograft) used for several types of tissue repair (e.g., nerve tissue repair) and evaluates the immunologic response and the degree of immunologic rejection after surgery.

The specification also supports the amendments to the claims 41 and 42, specifically paragraph [0011], which defines the basal laminae and endoneurium layer retain substantially the native extracellular

matrix structure as retaining the natural and generally original structure of the basal laminae and endoneurium layer. The cellular components are specifically removed without significant alteration of the natural extracellular structure of the native extracellular matrix (ECM). The structure is preserved (referred to as intact structural components), specifically, the basal laminae and endoneurium/endothelial layer retain their natural and generally original structure. In addition, paragraph [0036] defines the removal of cells without creating structural damage (thereby retaining extracellular matrix and essential components).

Claim Rejections - Claims 1-4, 7, 9-19 and 41-44 are rejected under 35 U.S.C. § 112.

The Action rejects claims 1-4, 7, 9-19 and 41-44 based on not complying with the written description requirement of 35 U.S.C. § 112. The Action contends that the Triton X-200 listed in the specification provides examples of non-ionic detergents. Applicants assert that the specification provides examples of anionic detergents including Triton X-200.

The specification as filed (page 12, paragraph [0042]) provides examples of anionic detergents including Triton X-200. Triton X-200 is in-fact an anionic detergent. The manufacturer's product information sheet (attached as Appendix A and incorporated herein) lists Triton X-200 as an anionic detergent. Similarly, the Sigma-Aldrich detergent product index (attached as Appendix B and incorporated herein) lists Triton X-200 as an anionic detergent. Therefore, the specification does provide specific examples of anionic detergents. These are described in a way that the skilled artisan would know the inventors had possession of the claimed invention and fully complied with 35 U.S.C. § 112 first paragraph. The claims also particularly point out and distinctly claim the invention and fully complied with 35 U.S.C. § 112 second paragraph.

As such, the specification satisfies the written description requirement under 35 U.S.C. § 112. For the reasons mentioned above, the Applicants respectfully request the withdrawal of the rejection under 35 U.S.C. § 112.

Claims rejected under 35 U.S.C. § 102(b) as being anticipated.

The structure, properties and characteristics of the product of the present invention are very different from the structure, properties and characteristics of the products disclosed in Livesey, Dennis, Gulati, Tanagho and Atala. When assessing the patentability of product-by-process claims over the prior art, the structure implied by the process steps must be considered, especially where the product can be defined by the process steps by which the product is made, or where the manufacturing process steps would be expected to impart distinctive structural characteristics to the final product. See, e.g., In re-

Garnero, 412 F.2d 276, 279, 162 USPQ 221, 223 (CCPA 1979). The products of claim 1-4, 7, 9-19 and 41-44 can be defined by the process steps by which the products are made and the process steps create distinctive structural characteristics in the final products.

The steps and materials used to prepare the graft of the present invention and the grafts of the cited references are different in both structure and physical characteristics and as a result, each of the final products are different in both structure and physical characteristics. For example, the attached (see Appendix C and incorporated by reference herein) Tissue Engineering article pages 1641-1651 (Volume 10, Number 11/12, 2004) (hereafter referred to as "Hudson") illustrates the importance of maintaining the internal structure and extracellular matrix components of a nerve tissue graft and compares the different methods of manufacturing nerve grafts. Hudson compares (page 1642 methods and materials section) a nerve tissue graft made by treatment with SB-10 (referred to in Hudson as "OA"), a chemical treatment method using sodium deoxycholate similar to United States Patent Number 6,371,992 (referred to in Hudson as "Sondell") and a freeze thay method (referred to in Hudson as "F-T").

Hudson provides in Figure 5 (page 1647) an image of the cross-sections of basal laminae visualized by laminin staining that compares the basal laminae after treatment with SB-10 (referred to in Hudson as "OA"), a chemical treatment method using sodium deoxycholate similar to United States Patent Number 6,371,992 (referred to in Hudson as "Sondell") and a freeze thaw method (referred to in Hudson as "F-T"). The different treatments produce different products with different structures and different treatments; Figure 5 of Hudson shows the cross sections of the basal laminae (i.e., the rings) after treatment, illustrating the fresh nerve tissue (Figure 5a of Hudson) and the SB-10 treated nerve tissue (Figure 5b of Hudson) have intact basal laminaes. In contrast, the chemical treatment with sodium deoxycholate by Sondell disrupted the basal laminae (Figure 5d of Hudson).

The different treatments produce different products having different characteristics. For example, Figure 7 of Hudson is a graph that compares the capacity to support regeneration or the axon density at both 28 and 84 days, in a fresh graft, in an OA treated graft (i.e., the sample with SB-10), in a graft treated with sodium deoxycholate (as in Sondell) and in a graft treated by the freeze thaw method. Figure 7 of Hudson shows the highest axon density at both 28 and 84 days is seen in the OA treated graft (i.e., the sample with SB-10). A decrease in the axon density from 28 days to 84 days is seen for the sodium deoxycholate treated graft (as in Sondell) and the freeze thaw graft (F/T). Hudson states that the OA graft showed a 910% higher axon density compared to the freeze thaw graft and a 401% higher axon density compared to the sodium deoxycholate treated graft (Sondell) (page 1650).

The treatment with SB-10 (e.g., OA treated graft of Hudson) showed a high capacity to support

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regeneration and maintained the extracellular matrix components and structure. In contrast, the graft prepared by the freeze thaw method showed a lower capacity to support regeneration and did not remove the cellular debris; similarly, the graft prepared by the sodium deoxycholate treatment (as in Sondell) showed a lower capacity to support regeneration and did not retain the extracellular matrix and structure (see page 1648 and Figure 7 of Hudson). In addition, the treatment with SB-10 (e.g., the OA treated graft of Hudson) did not show an increase in the CD8+ cells, and indicated that a rejection reaction was not present (page 1649 of Hudson). The process steps by which the products are made impart distinctive structures. characteristics and properties to the final product.

Therefore, products made by different methods (e.g., freeze thaw treatments, chemical treatments and the present invention) having different steps, mechanisms and reagents impart different characteristics and properties on their respective products. Thus, the products made by these processes cannot be identical products.

Claims 15-17 and 19 are rejected under 35 U.S.C. § 102(b) as being anticipated by Livesey, et al.

Applicants disagree with the Action's analysis of U.S. Patent No. 5,336,616 to Livesey, et al., ("Livesey"), which is said to disclose the claimed invention. Livesey does not anticipate claims 15-17 and 19 of the present invention. Livesey does not disclose the limitations related to nerve tissue, structural integrity, the reduction in the immune response of the graft or the enhanced capacity for regeneration produced by the present invention. As such, Livesey simply cannot anticipate the present invention.

The process steps of the present invention and Livesey are different; and, as a result, the products defined by the processes are different. Livesey discloses a product that is made by a process using different chemical agents to produce a product that has different properties than the product of the present invention. The chemical agents disclosed by Livesev include Triton X-100, polyoxyethylene (20) sorbitan mono-oleate and polyoxyethylene (80) sorbitan mono-oleate (Tween 20 and 80) (c.9, 11.41-52) which are non-ionic and sodium deoxycholate, and sodium dodecyl sulfate (id.) which are anionic. After Livesey decellularizes the tissue, it is incubated in a cryopreservation solution and cryopreserved. Furthermore, Livesey does not disclose sulfobetaines alone or in combination with an anionic surfaceactive detergent.

Although chemicals can be lumped into broad categories of generally similar characteristics, it cannot be said that each individual chemical of that category is the same (e.g., all organic compounds can Reply to Office Action of Sept. 18, 2006

be placed into one category-organics, but all organic compounds are not the same). Similarly, detergents are different characteristics (e.g., composition, structure, characteristics, charge, size, etc.) and as such their interaction with a substrate is different because it is dictated by the characteristics of the detergent. Although, some general characteristics of being a detergent may be shared, the individual compounds are not interchangeable as their specific compositions, structures, characteristics, charges and sizes are different

The chemicals used in the present invention and Livesev have different structures, different chemical formulas and different characteristics; therefore, the products produced by these processes cannot be identical. The skilled artisan knows that different compounds have different properties (e.g., the critical micelle concentration value, solubility, amount of damage to protein structures, amount of myelin basic protein removed and so forth) and with different properties, the ability to decellularize tissue is different. As the degree of decellularization is different, the product must be different. With differences in the degree of decellularization, the present invention and the product of Livesey cannot be identical. Therefore, the process of the present invention imparts distinctive structural characteristics to the final product.

Applicants respectfully submit that the Livesey fails to meet the standard of 35 U.S.C. § 102(b). As such, Livesey does not anticipate any of the claims of the present invention. Applicants respectfully request the withdrawal of the rejection under 35 U.S.C. §102(b).

Claims 15-19 are rejected under 35 U.S.C. § 102(b) as being anticipated by Dennis, et al.

The Action rejects claims 15-19 under 35 U.S.C. § 102(b) as anticipated by Dennis, et al., ("Dennis") (U.S. Patent No. 6,207,451), which is said to disclose the claimed invention. Applicants respectfully submit that the cited reference fails to meet the standard of 35 U.S.C. § 102(b).

The products of claim 15-19 can be defined by the process steps by which the products are made and these process steps impart distinctive structural characteristics to the final product. The process steps of the present invention and Dennis are different and, as a result, the products defined by the processes are different. Dennis teaches acellularized muscle anchors made by removing the muscles tissue from a subject, cut the muscles tissue into strips and pinned them to a substrate. The muscle strips are treated with a NaN₃ solution a deoxycholic acid (sodium salt), a solution of SDS and a solution of TRITON X-100

Dennis teaches a product made by a process using mammalian muscle construct, which is developed in vitro from cells extracted from mammals. First, Dennis relates to muscle tissue. Second, the product made by the process of Dennis and the product made by the present invention are different. Third, Dennis does not disclose the treatment with one or more sulfobetaines, nor does Dennis disclose the treatment with sulfobetaines and an anionic surface-active detergent. Dennis and the present invention use very different processes and reagents. As a result these differences impart distinctive structural characteristics to the respective final products (for the same reason as stated above).

Applicants respectfully submit that Dennis fails to meet the standard of 35 U.S.C. § 102(b). The products of the present invention can be defined by the process steps by which they are made and those process steps impart distinctive structural characteristics to the final products. As such Dennis does not anticipate any of the claims of the present invention. Applicants respectfully request the withdrawal of the rejection under 35 U.S.C. §102(b).

Claims 15-19 are rejected under 35 U.S.C. § 102(b) as being anticipated by Gulati, et al.

The Action rejects claims 15-19 under 35 U.S.C. § 102(b) as anticipated by Gulati, et al., ("Gulati"), which is said to disclose the claimed invention. Applicants respectfully submit that the cited reference fails to meet the standard of 35 U.S.C. § 102(b).

The products of claim 15-19 of the present invention can be defined by the process steps by which the products are made and the process steps impart distinctive structural characteristics to the final product. The process steps of the present invention and Gulati are different; and, as a result, the products defined by the processes are different. Gulati discloses a product that is made by a process of harvesting degenerated nerve cells and repeatedly freezing them in N₂ (I). Gulati then places the nerve cell on a dish of cultured cells (see page 120, section 2.3). It is unclear how a single nerve cell on an in vitro tissue culture that is repeatedly frozen and thawed is the same as a native, cell-free tissue replacement. Regardless, the process of Gulati creates a product that has a different composition, structure and characteristics than the product of the present invention.

Gulati does not disclose a tissue replacement made by soaking a tissue in a solution having one or more sulfobetaines, washing the tissue replacement in one or more solutions of a buffered salt, extracting with an anionic surface-active detergent and washing the tissue replacement in one or more solutions of a buffered salt. Gulati and the present invention are clearly different in processes and as such impart distinctive structural characteristics to the final product. Therefore, the product in Gulati and the product of the present invention are different, made by different processes and possess different characteristics.

Applicants respectfully submit that the Gulati fails to meet the standard of 35 U.S.C. § 102(b). As such Gulati does not anticipate any of the claims of the present invention. Applicants respectfully

request the withdrawal of the rejection under 35 U.S.C. §102(b).

Claims 15-19 are rejected under 35 U.S.C. § 102(e) as being anticipated by Tanagho, et al.

The Action rejects claims 15-19 under 35 U.S.C. § 102(b) as anticipated by Tanagho, et al., United States Patent Number 6,371,992 ("Tanagho"), which is said to disclose the claimed invention. Applicants respectfully submit that the cited reference fails to meet the standard of 35 U.S.C. § 102(e).

The process steps of the present invention and the process steps in Tanagho are different; and, as a result, the products made by these processes are different. Tanagho discloses a product that is made using a chemical treatment that includes a sodium deoxycholate solution containing sodium azide to remove cell membranes and intracellular lipids from the intermediate matrix. Tanagho <u>does not</u> disclose a nerve tissue replacement product obtained by a soaking an obtained nerve tissue replacement in a solution having one or more sulfobetaines, washing the tissue replacement in one or more solutions of a buffered salt, extracting with an anionic surface-active detergent and washing the tissue replacement in one or more solutions of a buffered salt.

The skilled artisan knows that different compounds have different properties (e.g., the critical micelle concentration value, solubility, amount of damage to protein structures, amount of myelin basic protein removed and so forth) and with different properties, the ability to decellularize tissue is different. As the degree of decellularization is different, the final product must be different, as shown in Figure 10. Therefore, the differences in the degree of decellularization between the product of the present invention and the product of Tanagho result in the products being different (e.g., having different compositions, structures and characteristics). The different properties of the compounds used in Tanagho and the present invention result in different components being removed from the tissue to form a product having a unique internal structure with different extracellular matrix (ECM) components. The process of the present invention imparts distinctive structural characteristics to the final product. Thus, the product of Tanagho does not maintain the same composition as the product of the present invention so they cannot be identical.

Applicants respectfully submit that the Tanagho fails to meet the standard of 35 U.S.C. § 102(e).

As such Tanagho does not anticipate any of the claims of the present invention. Applicants respectfully request the withdrawal of the rejection under 35 U.S.C. §102(e).

Claim Rejections - Claims 15-19 are rejected under 35 U.S.C. § 102(e) as being anticipated by Atala.

The Action rejects claims 15-19 under 35 U.S.C. § 102(e) as anticipated by Atala, United States

Patent Number 6,376,244 ("Atala"), which is said to disclose the claimed invention. Applicants

respectfully submit that the cited reference fails to meet the standard of 35 U.S.C. § 102(e).

First, Atala relates to an organ or part of an organ. Second, the process steps of the present invention and Atala are different and as a result, the products defined by those processes are different. Third, Atala discloses a product that is made using severe mechanical treatments using a magnetic stir plate and a paddle or a rotator platform. In contrast, the present invention provides a tissue replacement product obtained by soaking an obtained tissue replacement in a solution having one or more sulfobetaines, washing the tissue replacement in one or more solutions of a buffered salt, extracting with an anionic surface-active detergent and washing the tissue replacement in one or more solutions of a buffered salt. The process used in Atala and the present invention are different and impart different characteristics on the respective products. These distinctly different processes result in distinctly different final products. Thus, the product of Atala does not maintain the same composition as the product of the present invention so they cannot be identical.

Applicants respectfully submit that the Atala fails to meet the standard of 35 U.S.C. § 102(e). As such, Atala does not anticipate any of the claims of the present invention. Applicants respectfully request the withdrawal of the rejection under 35 U.S.C. § 102(e).

Claims 1-3, 9-14 and 17 are rejected under 35 U.S.C. § 103 as being unpatentable over Livesey in view of "Detergent Properties and Applications"

Applicants respectfully submit that claims 1-3, 9-14 and 17 are not obvious over the cited art and are, therefore, allowable under 35 U.S.C. § 103(a) for the reasons stated below.

A prima facie case of obviousness has not been established as (1) the prior art or combined references does not teach or suggest all the claim limitations, (2) there is no reasonable expectation of success and (3) there is no suggestion or motivation in the prior art to modify the reference or to combine reference teachings as proposed.

The Action states it would have been obvious to combine Livesey with a reference entitled,
"Detergent Properties and Applications" to achieve the present invention. Livesey as discussed, supra
(arguments incorporated herein by reference) does not include each and every limitation of the present
invention. Livesey does not disclose nerve tissue replacements, does not disclose treatment with
sulfobetaines, and the product formed by Livesey is different than the product of the present invention.
The "Detergent Properties and Applications" reference is merely a list lumping the zwitterionic detergents
together; however, each of the detergents are different, each having different structures, characteristics
and properties. The addition of the cited reference does not cure the deficiencies of Livesey, and even if
the cited reference did (which it does not), a prima facic case of obviousness would still not established

because there is not a reasonable expectation of success and no suggestion or motivation in the prior art to modify the reference or to combine reference teachings as proposed.

In addition, the Action's statement that the use of de-ionized distilled water would have been obvious is incorrect. Livesey taught the use of de-ionized water to wash off the fascia. In contrast, distilled water loosens the myelin sheaths (which are about 90% lipid) that surround the axons and swells in the presence of distilled water and allows the subsequent detergent solutions to penetrate and disrupt the cellular membranes of the myelin sheaths.

Accordingly, Applicants respectfully submit that the claims are not obvious over Livesey and the Sigma-Aldrich reference "Detergent Properties and Applications" and are, therefore, allowable under 35 U.S.C. § 103(a). Applicants respectfully request that the rejection of the claims be withdrawn.

Claim Rejections – Claims 4, 7 and 18 are rejected under 35 U.S.C. § 103 as being unpatentable over Livesey in view of Atala

Applicants respectfully submit that claims 4, 7 and 18 are not obvious over the cited art and are, therefore, allowable under 35 U.S.C. § 103(a) for the reasons stated below.

Neither Livesey or Atala (each of which are discussed *supra* and arguments incorporate herein by reference) nor any combination thereof teach or suggest all the claim limitations. Furthermore, there is no reasonable expectation of success and there is no suggestion or motivation in the prior art to modify the reference or to combine reference teachings as proposed. As such, a prima facie case of obviousness has not been established. Applicants respectfully request that the rejection of claims 4, 7 and 18 be withdrawn.

Conclusion

In light of the remarks and arguments presented above, Applicants respectfully submit that the claims in the Application are in condition for allowance. Favorable consideration and allowance of the pending claims 1-4, 7, 9-19 and 41-44 are therefore respectfully requested.

Applicants believe no fees are due at this time. If the Examiner has any questions or comments, or if further clarification is required, it is requested that the Examiner contact the undersigned at the telephone number listed below.

Dated: July 11, 2007.

Respectfully submitted, Chan I Lund

Chainey P. Singleton Reg. No. 53,598

ATTORNEY FOR APPLICANTS

Customer No. 34,725 Chalker Flores, LLP 2711 LBJ Freeway Suite 1036 Dallas, TX 75234 214.866.0001 Telephone 214.866.0010 Facsimile

Appendix A

Product Information

DOW Surfactants



Page 1 of 2

TRITON* X-200 Surfactant

Benefit			Applications	
Excellent detergent with tathering properties High stable foam Low toxicity Effective in hard water Good stability to electrolytes & chlorine		Personal care applications Emulsion polymerization Mild alkaline cleaners Heavy duty cleaners		
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	si Properties		Performance Properties	
Actives, v	M5%	28	Equilibrium surface tension', dynes/cm	30
Solvent Appearar	oce ·	Water Opaque, white iquid	Critical micelle concentration in distilled water at 25°C (77°F), ppm	970
pH, 5% a	noithles p	6.6	Draves 25 sec wetling conc. wt% at 25°C (77°F)	0.07
Viscosity	et 25°C (77°F), cP	7000	Ross-Miles Form Test, Initial/5 min.	
	125°C (77°F), g/mL	1.068	0.1% at 25°C (77°F), mm	88/81
	Closed Cup, ASTM D93	None -1 (30)	50°C (122°F)	155/7
	icional physical and chemical prop product Metarial Safety Date Shee		*Measured at 0.1 wi% and 25 °C (77 °F)	
Solubil	ity and Compatibility		Chemical Description	
 Solu 	ble in water		Name: Polyether sulfonate	
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Additional product information and performance data is available by requesting datasheets that are listed on the backside of this page.

Appendix A

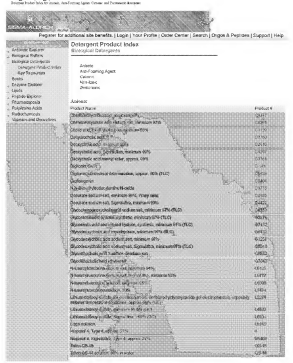
DOW Surfactants Page 2 of 2

Datasheets

- DOW Specialty Surfactants Reference Chart, 119-01491
- TRITON & TERGITOL Surfactants for Household, industrial & Institutional Cleaning CD, 119-01485-0501
- TRITON & TERGITOL Surfactants for Paint, Coatings, Adhesives, Stabilizers & Emulsion Polymerization CD, 119-01536
- Contact DOW Customer Service for current listing on conformance of TRITON Surfactants with U. S. FDA Regulations

Appendix B

Copy of information on the Sigma-Aldrich website listing of products classified by properties of detergents, e.g., Anionic and non-ionic.



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Appendix B

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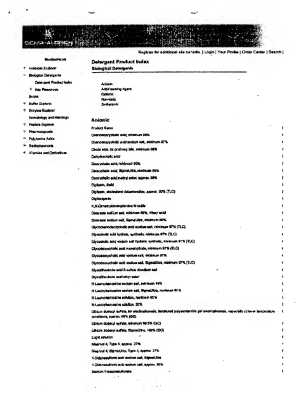
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 $Appendix \ B$ Descriptor Product index the Assistant, Arts-Fronting, Agradic Constant, and Zerithelineis desagnate

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Appendix B

Copy of information on the Sigma-Aldrich website listing of products classified by properties of detergents. INCLUDING THE MISS CHARACTERIZATION OF TRITON X200.



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Appendix 6	
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Triton X-700 solution, 76% in water (dispersion)	2
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Tribut® X-100, Stansal Stra	
Trigget X-100; Perceptor and colors of three	3
Tribut® X-114	3
Fifter® X.186 salution, 70% in water	;
Tribot® X-30t advason, 70% in wedge	4
Tritor® XAGS nonzon, 70% in water	2
Fritan® X-4S	ş
Frien® X-705-70 settion, 70% or water	1
TWEEK® 20, VINDOM FORE	Í
TNEEN* 20, Low-penerida; Low-perbonyls	ŧ
TWEEN® 20. Signal Hra	ŧ
THE SH 20, Low-passass; Low-co-fornys	ŧ
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TMERN* 80	3
TWEEN® SQ Vincoln Hydid, Low Peroride	\$
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TWEEN® 80 solution, Liquid	ŧ
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Appendix B

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Appendix C

TISSUE ENGINEERING Volume 10, Number 11/12, 2004 © Mary Ann Liebert, Inc.

OPTIMIZED ACELLULAR NERVE GRAFT IS IMMUNOLOGICALLY TOLERATED AND SUPPORTS REGENERATION

Terry W. Hudson, Ph.D., Scott Zawko, B.S., Curt Deister, B.S., Scott Lundy, Char Y. Hu, Kate Lee, and Christine E. Schmidt, Ph.D.

Appendix C

TISSUE ENGINEERING Volume 10, Number 11/12, 2004 D Mary Ann Liebert, Inc.

Optimized Acellular Nerve Graft Is Immunologically Tolerated and Supports Regeneration

TERRY W. HUDSON, Ph.D., I SCOTT ZAWKO, B.S., I CURT DEISTER, B.S., I SCOTT LUNDY,2 CHAR V. HIL3 KATE LEE,3 and CHRISTINE E. SCHMIDT, Ph.D. 1.2.4

ABSTRACT

To replace the autologous graft as a clinical treatment of peripheral nerve injuries we developed an optimized acellular (OA) nerve graft that retains the extracellular structure of peripheral perve fissue via an improved chemical decellularization treatment. The process removes cellular membranes from tissue, thus efininating the antigens responsible for allograft rejection. In the present study, the immunogenicity and regenerative capacity of the OA grafts were tested. Histological examination of the levels of CD8* cells and macrophages that infiltrated the OA grafts suggested that the decellularization process averted cell-mediated rejection of the grafts, in a subsequent experiment, reponeration in OA grafts was communed with that in isografts (comparable to the clinical autograft) and two published accilular graft models. After 84 days, the axon density at the midpoints of OA grafts was statistically indistinguishable from that in isografts, 910% higher than in the thermally decellularized model described by Gulati (I. Neurosurg, 68, 117, 1988), and 461% higher than in the chemically decellularized model described by Sondell et al. (Brain Res. 795, 44, 1998). In summary, the results imply that OA grafts are immunologically tolerated and that the removal of reiislar material and preservation of the matrix are beneficial for promoting regeneration through an accitotar nerve graft.

INTRODUCTION

on stationary of severed peripheral nerves are treated Leither by surgical realignment of the individual nerve fascicles (i.e., primary neurocritaphy) or by implantation of an autologous perve graft (i.e., an autograft). Primary neurombanhy is performed if the nerve ends can be sutured together without inducing tension; otherwise, as autograft is typically used to bridge the cap between the severed nerve ends. Development of an equality effective replacement for the autograft is needed because the proorders entails multiple surgeries and the loss of function or sensation at the donor site 1.2 To date, no alternative he removed from donor tissue by several techniques in-

as effective as the automost at stimulating repeneration over long distances has been demonstrated.

The internel structure and extracellular matrix (ECM) components of a nerve graft have been shown to be critiest for guiding cell migration and nerve fiber ciongation.4-4 Thus, development of an acciliate agree graft, which contains the parend ECM components and structime but not make odly, could be valuable as an altermative to the current autograft. Accibing nerve grafts can also be used to study the roles of the ECM and cellular comments concornitantly

To crease acultular grafts, the reliable components can

^{*}Describes: of Chemical Passinerine, *Generoment of Biomedical Engineering, *Denotings of Biogram Reproduction, and *Texas Meterials Institute, University of Texas at Austin, Austin, Texas.

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chaling thermal? and chemical* processes. Thermal decellularization is the most common process in the litterature; It estable subjecting the tissue to repeate breate-thaw cycles. Although thermal decellularization does kill the cells and reader the guid generally nonimmunagenes? ³⁰ the process does not extract the cell menmants. As a practic, on estimate number of Schwaiss cells and materiphages invade the basal laminate pixels to learn in cellular televis during the first days after implantation. This cellular invasion proentially delays the regenerative process and disaugest the basal laminat. ³¹

Several chemical mentionats have been designed to reach energy gifts anothermonopine white also removing much of the cellular debris. However, chemical treatments cause more damage to the ECM thus therements cause more damage to the ECM thus theremate a collularization. 3,45 One of the most common observation decellularization protectols in the theretare was engineally developed by Johnson et al., 16 and later modified by Sondiell et al. 3

We previously developed a chemical decellularization oracess to create aptimized acaitular (OA) nerve spalits with an extracellular environment similar so that of natime nerve tissue, but without the cullular material that is believed to elicit cell-mediated rejection. 19 Thorough histological evidence was presented in thus article demonstrating both cril (e.g., Schwann cells) removal and ECM (e.g., basal lamines) preservation. In the present study we had two primary mosts; (1) so determine whether the removal of cellular components accomplished with the OA protocol translated into an intensaciogically scienared graft, and (2) to compare the regenerative capacity of the OA graft with that of other established scellular graft anodels Addressing the first goal of this week, we implanted CA gradis into rues of a different strain than the donor noimals (i.e., pllografts). Rejection was evaluated after 28 days on the basis of the level of immuse cells (e.g., T cells and macrophages) in the graft. 18 The level of cells expressing CDS' exsisties (i.e., sell surface markers on execucic T cells) and macrophage cells in the graits after 28 days demonstrated that the GA profits were not undergoing zeil-mediated rejection. Thus, the removal of cellular material translated into an immunolossically tolerated spaft.

Regeneration in OA grafts after 28 and 54 days was companed with been in every agrifts counsed according to published thermal and chomical describation for published thermal and chomical describation proportions. OF the other models were used to study the traportions of ECM preservation and cellular semonal in expensation capacity. Actor describative was significantly higher in OA grafts than in the other accellular models implying that preservation of the natural ECM and removal of cellular material are beneficial for regeneration through an acciliant nerve graft Even through required in the result of ECM and the contract that the contract the contract through an acciliant nerve graft Even through required in the result of the contract through the contract of the contract through the contract of the contract through an acciliant nerve graft Even through regions and the contract through the contract

to bead analy. This was necessary so that our data could be compared with published work on other auctiniar graft models that also employed the Ki-mm gap.

APACAMAN / P. B. A. CAMI

MATERIALS AND METHODS

Creation of grafts

To create OA grafts, been the left and right sessitic nerves were harvested under anoptic conditions from 350g Harlen Sprague-Dawley (HSD) male sots. The tissue was handled only on the ends to minimize structural damage. On harvest, the nerves were immediately placed in RPM1 1640 solution at 4°C. All subsequent steps were conducted in a luminar flow bood for signifity. Fatty and connective treate was removed from the nerve epirgenrium. The nerve tissue was cut into 15-mm segments and placed at a 15-mil conteal take filled with delenated distilled waste. All washing steps were curred out at 25°C with agitation. After 7 h, the water was aspirated and replaced by a solution containing 125 mM sulfaberaine-10 (SB-10), 10 mM obserbate, and 50 mM sedium. The nerves were agitated for 15 h. The tisaue was then desend for 15 min in a washing solution of 50 mM phosphate and 100 mM sedium. Next, the washing solution was replaced by a solution containing 0.14% Tritos X-266, 0.6 mM sulfobetaine-16 (SB-16), 10 mM phosphate, and 50 mM sodium. After agitation for 24 h. the tissue was rissed with the washing solution three times (5 min user ringe). The nerve segments were again agetated in the SB-10 solution (7 h), washed more, and agitated in the SB-16/Friton X-200 solution (15 to) Finaily, the tissue segments were washed three times (15 Sau stadgeodd Mea Of Snieithaus aodaioa an fragailte 50 atM andium and stored in the same mintion at 4°C.

Other acellades nervo gasts models were created according a published materials on a basis for companies. The chemically decellatanced model was created by a promoted published by Soudella et and Bristley, because the promoted published by Soudella et and Bristley, because the existse was agistated in distilled waters for 7 h. in 46 n in 97 thou, 7-100 is desulted water evernight, and then the 7 thou, 7-100 is desulted water evernight, and then in 98 nth solitum decoydolustics in distilled water for 26 h. These sceps were repended selence performing a final wash in distilled water. All mediators caps were performed as commiscingeration, and the fission was subsequently and in 10 nth pinaphun-buffered saline (PBS) solution at 4°C.

The internally decidinated model (i.e., a frenze-theoly graft) was constant functioning to the protocol decarbody by Galati. "I tensectionly after increes, nerve tissue was disposed in figuid increege for 60 s, and then for 1985 at room temperature for 60 s, and then the process was recommended from the first increed the form of the frenze-theory for 60 s, and then the process was recommended from admittenant times. The frenze-theory first grafts were placed in PBS at room semporature and used within 30 mile.

All chemicals were purchased from Sigma (St. Lauis, MO) unless otherwise noted. All solutions were suroslaved or filter wastlisted believe tree.

Implantation of grafts

bougarts and altografts were used to test the immunogenicity of the OA grafts. Isografts, which minute the autograft, were harvested from a donor minus! fe.g., Lewis rut) of the same strain as the bost agintal (e.g., Lewis rat). This served as a measure control for any manufacture sponse that results from the sergical procedure alone. Allografts were harvested from a donor animal (e.g., HSD rate of a different strain than the host anisted torg. Lewis rate. The differences between these rat strains are addressed in the Discussion (below). The fresh allograft served as a positive contool because it is known to elicit cell-mediated rejection. The OA isograft was used to exsmine the in visco response to the treatment protocol (r.g., response to residual chemicals). As OA allowed was inspected for residual antigens following our descitatarization accordant. The four experimental aundations tested are summarited in Table 1.

Each rat was anosthetized with an immperisonced injection of ketamine (120 mg/kg body weight; Webster Veterinary Supply, Sterling, MA) and xylazine (15 mg/kg body weight: Webster Veterinary Supply). The sciatic nerve on the right side was exposed, transcetted, and 5 mm of nerve was sumpress. The easts of the graft were trimmed immediately before implantation to attain a clean-cut, 10-mm graft, The graft was unused to both the presented and distal sorve ends, using 10-0 varyl sotteres (Ribison, Somerville, NJ). The muscule was drawn back together with 5-6 cheomic gut satures (Ethicon), and the skin was closed with wound clips (BD Diagnostics, Sparks, MD), Surgical methods were performed in accordance with regulations established by the National Research Council in the Guide for the Care and Use of Laborators Assimals. 20

Immunogenicity of grafts evaluated by histology

Grafic representing all four experimental conditions were harvested 28 days after implantation, Each anistral

was presentiational, and the nerve girdt was emposed. Before harvesting, the graft was fixed for 1 min with ghararakin/yak-with parallermalakin/yak-with PBS. The witstale carers, was then transacted 5 is some show and the the graft, the distal and was marked with a stick, and the graft was placed in fixative at 4°C. Alter 30 min, the was was transferred in PBS and storted at 4°C sortil it was senbedded in paraller.

Histology was used to inspect the allografts for signs. of impuspological rejection. The tissue was dobyerased with graded alcohol solutions and a ylone, and then conbedded in paratha. Langitudinal sections of tissue, 7 µm thick, were cut with a merratome and captured on glass: slides. Immunostaining was performed with onto-CD8a (BE) Biosciences Phorosingen, San Diego, CA) and ontimacrophage (Chemicon International, Tameculu, CA) primary antibodies. Horseradish permidese (HRP)messed secondary unibodies, 3.3 diaminobenzidine (DAB) substrate (Vector Laboratories, Burlingamo, CA), and an easin consterstain were used to viscolize the invasting cells. The stained sections were visualized on an Olympus 1X20 (Olympus America, Melville, NY) inverted mecroscope, and the images were captured with an Operopics Macmiffire (Goleta, CA) digital color camera. Images of the stained tiesus acctions were combined in Assobe Photoshop to create a composite of the entire graft. Using Scien brage software (Scion, Predarick, MA), the percentage of area of the gruft covered with positively stained CDR+ cells and macrophages was determined.

Acellular graft models compared in vivo

To study the impact of refultar dobtic and structural procession on regeneration, three activities graft models were examined in view, OA grafts. Sondell grafts, and F-T grafts were created as described in Materials and Rethods. Firstle grafts are a mixing of the clinical autograft and were included in the experiments as a positive control. The OA grafts and Sondell grafts were prepared within 30 days of implicatation. The time between himself and the process and implication of the F-T grafts and frestly states was nover jonger than 30 title. Donor and flort animals were HSD rate.

TABLE 1. IMPLANTS TO EXAMPLE PROGREDLEGICAL TOLERANCE OF COPYROSIS GRAPPS

Grafi typi	Division nomina	Missa sarurin	Number of implants	daulycing requence to:
Fresh isoeraft	Lewis	Lewis	3	Surgical procedure (acquirer rector)
	HSD*	CESSS	3	Suggical promodure (negative custral)
Fresh allowruft	Lewis	BISD	5	Natural anapers (positive control)
(Sprintiged anotheler integrals	HSD	RISD	.5	Feramons protocul
Ontimised sorbider allogation	Lawis .	SESES	5	Residual astigned

^{*}Lowin rank are an inhered stenies (i.e., greater than 98% genetic horstogeneity),

*1810 rate are an exchang stenie, but the animals used were from a closed cultury.

Histological comparison of decellularized tissues

A comparison of the ECM structure in the accellular parish before implication was constanced by visibilities give board basinar. The grafts were prepared to selectly described, excluded, and cross-sectioned and cross-sectioned and articlamining primary sufficient processing described primary sufficient sufficient primary sufficient primary sufficient sufficient primary sufficient primary sufficient sufficient primary sufficient suffi

Regenerative capacity of grafts evaluated by histology

Grafts were harvested 28 and 84 days after insplantation (Table 2). The numbers of acreesed grafts for each time point are out the same because some animals were killed early due to insemnialisation, which is consistent with the unconstitution in MSD was observed by others. 3

To evaluate the regenerative potential of the three occljular graft modela, longitudinal tistue sections were stained for regenerated axons, using the RT97 anti-acurofilament primary antibody (Developmental Studies Hybridoma Bank), an HRP-conjugated secondary outbloody, and DAB. Subsequently, caoss-sections were out from the enideoint of the englis and stained for near thioments. The stained sections were visualized with a ×20 objective and images were captured with a digital carnera. A 20 × 16 em image was printed for each sample. The number of nerve fibers in each image was counted, and the area of nerve cable in the image was necessard. Because a poytion of the nerve cable had been removed by sectioning the distre longitudinally before taking cross-sections, the got bleer of stones in each nerve cable could not be determined. Instead, exon density was culculated by divictims the number of nervo filters by the area of the cable from which the count was taken. Select specimens were not used in the axen density analysis if less than 33% of the nerve valide commined after longitudinal sectioning. The number of samples analyzed for each graft and time point is reposted with the axon density data. Regions of connective tissue at the periphery of the graft,

Table 2 Implants to Evaluate the Bouleverstyk Capacity of Councies Alphabas Grafts

Herrienied (28 days)	Hornest 184 éty
9	6
6	5
6	4
9	6
	528 dinjo) 9

based on noophological evaluation, were excluded from the analysis.

Spainneal analysis

Analysis of variance (ANOVA) was performed to deserment the statistical significance of the differences between neurals. Specifically, as Γ is see was used to deternance whether the variability between data sate was equal or unequal. A Γ state was then used to deterative whether the difference between the averages of the lata with was statistically significant. A significance levice of $\rho < 6.05$ was used as the cutoff Γ is, ρ values are reported only for causes in which $\rho < 0.055$.

RESULTS

OA grafts use immunolavically telerated

To evaluate the immunological response by a host to GA grafts, four experimental conditions were tested with sciatic nerve graft implants (Table 1). By staining longitudinal sections of grafts for cytotoxic T cells and macrophages, the level of cell-mediated immune response was determined. Elevated levels of cytotoxic T cells are expected in tissues undergoing cell-mediated rejection and increased levels of macroplange sells see expected in rejected allowrafts, However, macrophages are also recreated during Walterian degeneration in clear debris and referse regrotrophic factors for regenerating nerves. At 28 days, both cell types could be soon throughout the full length of all grafts (Figs. 1 and 3). The inflatration of CD8" cells into fresh altograffs was higher thus into fresh isografts (p < 0.01) and OA grafts (a < 0.005) (Fig. 2). Meanwhile, the levels of CDS+ coils in OA issgrafts and OA allografts were lower than those observed in firesh isosesits (p < 0.05). Macrophage invasion isto fresh isografts was lower than into fresh altografts (p < 0.05), but the differences between other andre were not statistically significant (Fig. 4). Thus, histological engageration of the levels of CD8+ cells and reacrophages that infilterated OA grafts suggened that the decellularization process overest callenedistesi veiestion of the orafts.

OA process preserves the ECM

Images of tissue sections resented for luminor allowcontrol of heads luminor generation among the decellularization prosects (Fig. 5). The ringillies structure in native nerve issue are upon columns of best illuminor (Fig. 5a), and similar structures are apparent in tissue treated aeroming to the OA protects (Fig. 5b) and the F-T protect (Fig. 5c). The head luminous appear high fragmented in tissue created seconding to the Sondel's prosecut (Fig. 5c).

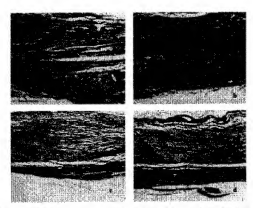


FIG. 1. Longitudinal sections or insers were our from tail fresh integrable, the first hallegraft, etc. DA. Insegnaft, and coll. CAS. Insegnaft, and coll. CAS. Insegnaft harmonic CAS days after integrationation. Those Sections were statisfied for CAS. In surface market on explosious T-CAS. In serial can destinate in the fresh altegrafts was estably higher, but the CAS grafts appeared indistinguishabit from the first hisografts. Seed here: 20% and the CAS grafts appeared indistinguishabit from the first hisografts.

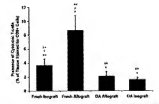
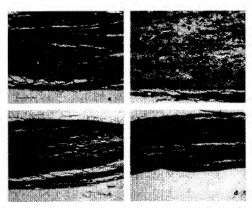


FIG. 2. Cell-medianed immune response in fresh and GA nerve grades was avaluated by determining the percentage of its outcovered by CDS* exils. Feels illuspoints demonstrated a satisfacility significant develope in CDS* code. GA records and offigards were subsidiesly indistinguishable from the fits superfits, indicating time cell-mediated immune experience was societized goody in fresh inlargation. Symbols above the columns designate in significant ciril forcine from fresh inequality. Special illuspoint (§ 7s. Code) singuisht (§). And Onlargatic (§ 4).



FKG. 3. Longishalinad seatonin- of tissue were two from (a) found inograths, (b) fresh tillograths, (c) OA inograths, and (d) OA altograths have seen 12 days after imploatation. These periods were shared for innoverplayer, its times colls involved in Waller has degenerated, nother pergeneration, and issue definition, and issue definition.

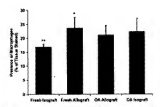
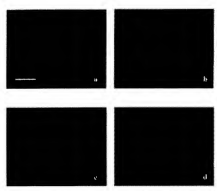


FIG. 4. Level of stateophingue present in Both and OA derive gifter short 25 days was reducted by determining the precentage of a text stateoid in Medical Indian Section 11 and 12 and



FRG. 5. Consecuções or best famises voco visualized by famisins salariag. The regifice appearance of the opin historia visia from several contraction of the opin historia for a large from several contraction. Of the best families for the opin historia for a large from the contraction of the best families families with distinguish for (d) a Sondell graft, suggesting that the basal lantaine were shreaged during the decellularization treatment. Scale but: 19 ann.

OA grafts support regenerating awas

The capacity of the OA graft to support nerve regeneration was tessed by examining the growth of axons through the various nerve isografts other 28 and 84 days. All she grafts were isografts, harvested from and implented to by 1800 rats. The grafts included (1) fresh isolated (2)

grafts, (2) OA grafts, (3) Sondell grafts, and (4) F-T grafts. Longitudinal sections and errors-sections of the grafts were stationed for neurofilaments (i.e. cytoxicalizad practical found in axons), As 28 days, new axons had grown completely axonse the grafts (Fig. 6). The axons anomared to meet ensistance receiping four the proximal



FIG. 6. Another aggreeration through 28-day OA series goalts was demonstrated by deliving longicalised disconsistents for internationation. Random patterns in the cases at the junctions of the (a) positional server and grait used set into grait and similar terror suggest a long displace as one amount of season at the control of the grait. However, some we (b) the indigitant of the grait work julying alaqued, suggesting that they were guided by the extraordisals retructure of the grait. Susteric marks (5) at the wave-grait amonthou see shown in (a) was (a). Such that (100 µm.

nerve and into the scalt and from the sraft into the distal nerve and, as demonstrated by the nordinearity of newrefilturenss around the sumre points (Fig. 6a and 66c). However, ourse the axons extended into the graft, they grew linearty, as demonstrated by the parallel neurofilemems at the raidpoint of the graft (Fig. 6b). Similarly, the axons grew linearly in the distal direction once they extended into the distal nerve and (data not shown). The some pattern was observed in the 84-day QA grafts. Histological stations on longitudinal sections also showed that Schwaan calls were present throughout all graft types at both time excises (data and shown). Thus, the OA nerve grafts supported axonal regeneration and golded axons toward the distal nerve and

Regenerative copacity of optimized graft suspasses other acethius models

In addition to visually examining the growth of axons through the grafts, exen density in grafts was determined. The same OA grafts and theth kografts that were harvested 28 and 84 days after implantation and sectioned longitudinally were subsequently cross-sectioned at the midpoint, stained for newofilements, and examined, in the 28-day grafts, the fresh grafts (a = 6) and OA statts (a = 7) were nearly idensical with auon densities of 0.9 and 6.98 acous/106 pm², suspectively (Fig. 7). The F-T grafts (n = 5) had 0.50 acous/100 µm2, and the Somicil grafts (n = 6) had 0.69 axons/160 μ m². Axon density in the F-T grafts was significantly lower than in the fresh grafts and the OA grafts (p < 0.01). Axon density in the case profesentially grow through the basel families tubes

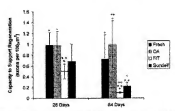
Sondell grafts was also significantly lower than in the fresh grafts (p < 0.01) and the GA grafts (p < 0.05).

Fresh grafts (a = 5) and OA grafts (a = 5) horvested after 84 days were still rost argusfavorily different, with axon dentities of 0.73 and 6.92 arons/100 acm2, respectively (Fig. 7). The F-T grades (a = 3) and 0.10 shores/100 μm^2 , and the Soudell grade in ≈ 3) had 0.23 axom/100 acm2. Axon density in the F-T grads was significantly Lower than in the fresh grafts (p < 0.05) and the OA grafts (p < 0.65). Axon density in the Soudell grafts was not significantly lower than in the fresh grafts, but was sigpifficantly lower than in the OA grafts (p < 0.95)

Because the freeze-shaw decellularization process joge not remove celistist debris and the Smaleli decella-Surkention process does not preserve the ECM, the highest grow densities as 24 and 84 days in the OA grafts suggest that removing cellular debriv and preserving the ECM improve the regenerative capacity of acellular nerve

DISCUSSION

An alternative method for treating severed peripheral nerves is asseded to avoid multiple surgenes, sonor sixe morbidity, and other drawbacks associated with the antograft. Accliniar nerve grafts, derived from donor nerve tissue, are composed of endogenous sissue proteins. Because of their natural composition and the fact that ax-



PRG. 7. The regenerative capacity of finer serve graft models was evaluated by recasoring area denvity in cross-sections of the and is 26 days after implantation and 84 days after implantation. From megrafic served at a social for the autograft (positive comhold Axon density in hesh grafts and OA grafts was standardly indistinguishable. P/T grafts had the howest more density, imphysic that the prevence of cellular Johns may reduce the regenerative capacity of an abeliake good. Southall grafts demonstrated a singuistically fower agon decreay that OA grads after 84 days, suggesting that preservation of the ECM introded the regenerative expecting of DA grafts. Symbols above the cotacins designists a significant difference from Stock graft (**), GA graft (**), FT (w), and Standard (++) graft.

found in nerve riscue, aceildar nerve grafts exhibit potential for use as a next-generation nerve graft. We bypothesized that improving the decellularization process to yield a bener-preserved ECM would lead to an invprovement in resentation. However, for OA grafts to be used clinically, they must also be immunologically tolerated. In previous work, we developed a ratthod of se-

WE ENTIRED A CHARLEST AND THE VALUE OF

moving the cellular material believed to be responsible for internativelegical adjection while also preserving the BCM of nerve tissue, in the current work, the OA graft was tested to vivo to determine its regenerative caracity and impunpmentative

Cellular actiones are eredominantly resounsible for the immunological rejection associated with nerve allografts, particularly the antigens associated with Schwana cells, endothelial cells, and macrophages, 18,22,23. The removal of cellular components by the OA protocol was correband with the iransprological response to an altograft. The major histocompatibility complex (MHC) of the rat is called RTs and is highly polysporphic. 24 Rat strains can be characterized by their RTI haplotype (e.g., RTIb), RTd, RTH). Matching of haplotypes plays a productionest role in allograft survival. Gulati and Cole demonstrated that in allografts involving strains of different RT1 haplotypes, the increased presence of transons cells associated with rejection was readily dejectable at 28 thrys. 68 Thus, their nerve sissue from an HSD ret (RTTh) implanted into a Lowis ret (RTH) (i.e., a fresh allograft) should display signs of impernological rejection after 28 days, Similarly, an aredfular altograft should be rejected if the graft commins membrase-bound entirens associated with the RTI haplotype.

Rag cytoloxia: T cells earry a CD8 cell surface marker fi.e., they are CDR* cells), and disconsence of evictorise T calls is an important indicator of cell-mediated graft rejections. However, a maximum number of CD8° cells that are not evolutionic should be present in any nerve graff after 28 days, whether or not it is undergoing rejection. The noncytosoxic CDS+ cells are a subset of macenphages that are known to invade after scintic nerve insuries, even in the absence of resection.25 Macrophages are increase cells that respond to serve injury. Melear celhator debris during nerve degeneration, 25 and support regeneration by inducing and producing growth factors.28 In the case of a sejected allograft, higher numbers of macreplaces should be present. 26 However, macrophases also restiond to other cues in the regenerating nerve, so an increase in macrophages without a concomitant increase in CDS+ cells dues not indicate miection. Thus, the presence of CDS* cells and macrophages was unticinsted in all four graft models. However, a statistically significant increase in both CD8* cells and macroplages in a graft, when compared with a tresh isograft, would indicate that the graft was undergoing cell-mediated asmution.

Iremanological tolerance of OA grafts was confirmed

As anticipated, the fresh allograms exhibited a watistical increase in both CD8* cells and macrophages compared with facali isografia (Figs. 2 and 4). The OA alloerafts did not show an increase in CD8* cells compared with fresh isografts, indicating that they did not eitest resection. Further evidence that the OA allografts were not rejected are the similar levels of CD87 cells and macrophages in the OA isografts and OA allografts.

Macrophage invasion into the OA grafts appeared pliebtly biober than invasion into the fresh isografix, although not signifugativ. A ransible cause for the elevated level of macrophages in the OA grafts compared with the fresh isografts is that the open based liming tabes and the absence of morein permitted a greater number of resurronisspes to specife and semain inside the OA grafts. This may be beneficial because macrophages produce argueth factors, in summary, the untigens that would have initiated or Benediated immunological rejection of OA allogisfis were removed.

Regenerative capacity correlated to graft structure and content

The two design criteria for the OA grafts were to remesve cellular maserial and to provide structural support for regenerating nerves, it was hyperhesized that this would improve regeneration in comparison with other accilcits grafts. The importance of structural support was revealed through histological examination of longitudinai risque sections. Axons grew linearty to regions of defixed structure (e.g., in the nerve graft and distal nerve cable), but their path was irregular in regions where the post was assuched to the nerve ends (Fig. 6). The irregular parteens were potentially caused by the misalignment of basel language at the junctions between the nerve ends and the graft. As the axons crossed into and out of the eraff, they had so find new basal laminac to provide them with guidance

to addition to providing epidence. OA grafts also supported higher axes densities after 14 and 84 days then did other published accliniar graft models (Fig. 7). The lowest axon densities were found in F-T grafts. Atthough the structural preservation in P-T grafts was similar to that in optimized grafts (Fig. 5), the F-T procedure was the only decellatarization procedure that did not remove cethalar debris. Thus, a correlation is suggested between the presence of coll debris and a reduction in the level of nerve regeneration. The primary of flerence between Sondell grads and OA grafts was preservation of the ECM (Fig. 5). Consequently, the higher axon slessity in OA gentles suggests that providing regenerating atoms with an RCM structure that minutes earlier serve is important for maximizing repeneration is an acellular graft. The un-

purtance of those factors appears to become more evident over lenger time periods, with the OA graft demonstrating accur densities 910% higher than the F-T graft and 401% higher than the founded graft after 84 days.

soft or eggeter tests the stronteur galax and or or acquilencame treats integrated were the only greets that proclaimed living cells (e.g., Schwarm cells and misconsplanes) religious associations that any or of the accidiant grafts. The data suggest first in 18-min nerve grafts, the combination of destatable extentive and the removal of cellshift debris was sufficient to attain acquidenties statistically indistinguishmide from these in fresh longarth (Fig. 7), the case of longer grafts, however, the rend for support sells is expected to be once careful. The OA graft can be to to reas lippires with longer gaps by incorporation of cells (e.g., Schwarm cells) left from implantation.

This work suspenses that the OA graft may serve as a starting template for an off-the-shelf nerve graft. In addition, this graft is well suited for studying specific aspects of nerve regeneration. Cellular components \$10.51 (e.g., Schwann cells and macrophages) and growth factors³² are important for successful peripheral nerve regeneration. Excellent research is being performed with growth factors in fabricated systems. 15-15 but the interaction of those components with the natural nerve environment is also insportant and could lead to further improvements. The natural structural environment of the OA graft maters is an ideal model for studying these interactions and for examining individual cell types and growth factors through soluctive incompration into the eraft. As more information is gained about the role of the ECM, support cells, and growth factors, better therapeutic systems can be engineered for stimulating nerve regeneration.

ACKNOWLEDGMENTS

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Address teoprist requests to: Dr. Cartelius Schmidt Department of Biomedical Engineering, MCFC0800 University of Texas et Austin Austin, TX 78712

E-moif: Schmidt@che.stexas.edu

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